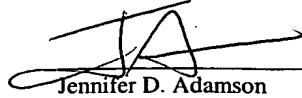


Patent
TS-1309 (US)
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Jennifer D. Adamson

Date: February 9, 2004

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re application of)	
)	
JAN K. F. BUIJINK ET AL)	
)	
Serial No. 10/720921)	Group Art Unit:
)	
Filed November 24, 2003)	Examiner:
)	
CATALYST PREPARATION)	February 9, 2004
)	

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CLAIM TO PRIORITY

Applicants reaffirm the claim for the benefit of filing date of the following foreign patent applications referred to in Applicants' Declaration:

EPC Application Serial No. 02258296.9 and 02258294.4 - filed December 2, 2002

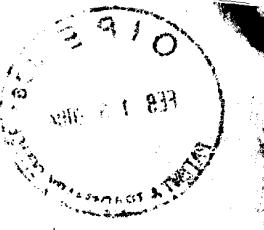
Copies of the applications certified by the European Patent Office are enclosed.

Respectfully submitted,
JAN K. F. BUIJINK ET AL

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Enclosures





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Attestation

Die angehefteten Unterlagen stimmen mit der ursprünglich eingereichten Fassung der auf dem nächsten Blatt bezeichneten europäischen Patentanmeldung überein.

The attached documents are exact copies of the European patent application described on the following page, as originally filed.

Les documents fixés à cette attestation sont conformes à la version initialement déposée de la demande de brevet européen spécifiée à la page suivante.

Patentanmeldung Nr. Patent application No. Demande de brevet n°

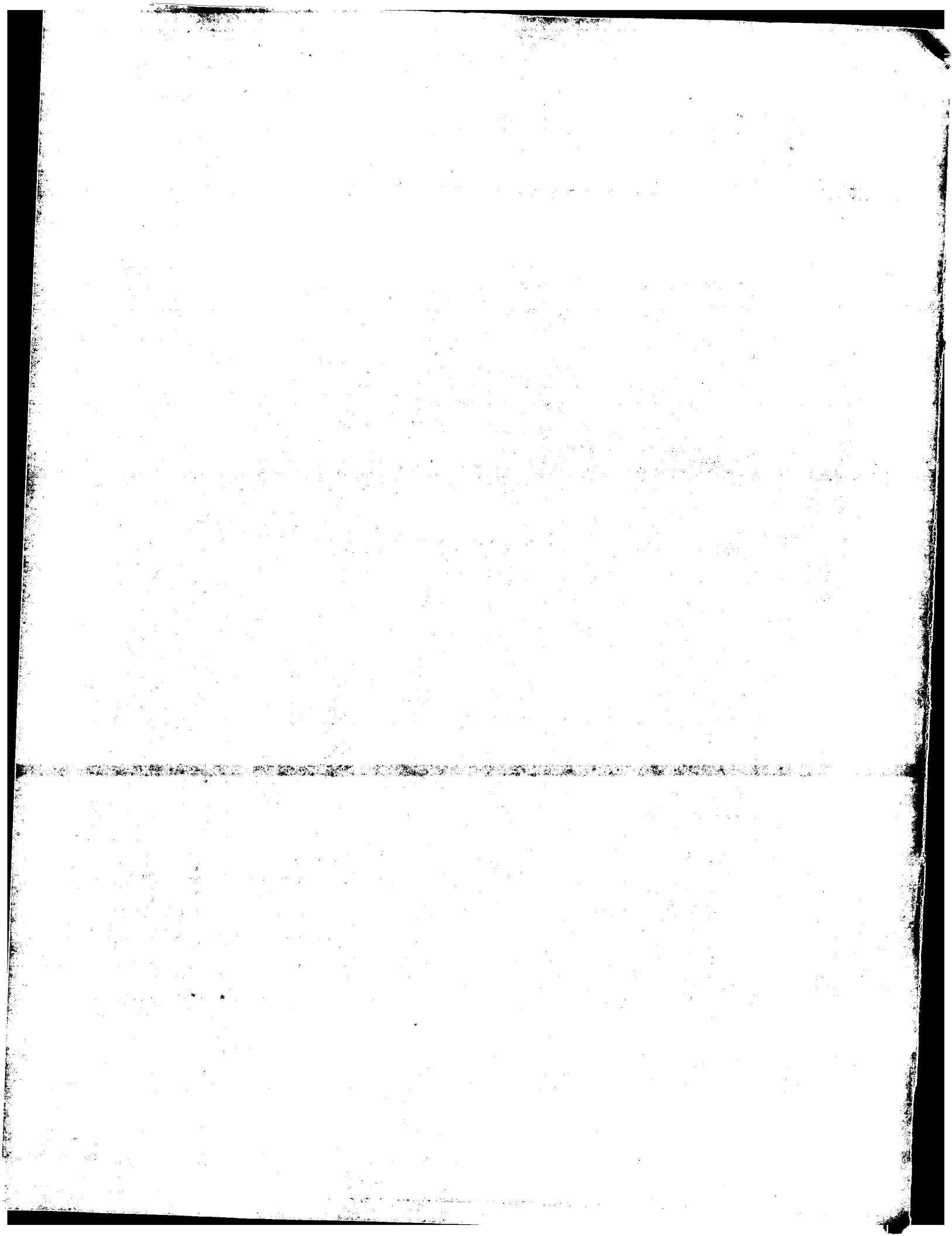
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Der Präsident des Europäischen Patentamts;
im Auftrag

For the President of the European Patent Office

Le Président de l'Office européen des brevets
p.o.

R C van Dijk





Anmeldung Nr:
Application no.: 02258296.9
Demande no:

Anmelde tag:
Date of filing: 02.12.02
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Anmelder/Applicant(s)/Demandeur(s):

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Bezeichnung der Erfindung/Title of the invention/Titre de l'invention:
(Falls die Bezeichnung der Erfindung nicht angegeben ist, siehe Beschreibung.
If no title is shown please refer to the description.
Si aucun titre n'est indiqué se referer à la description.)

Catalyst preparation

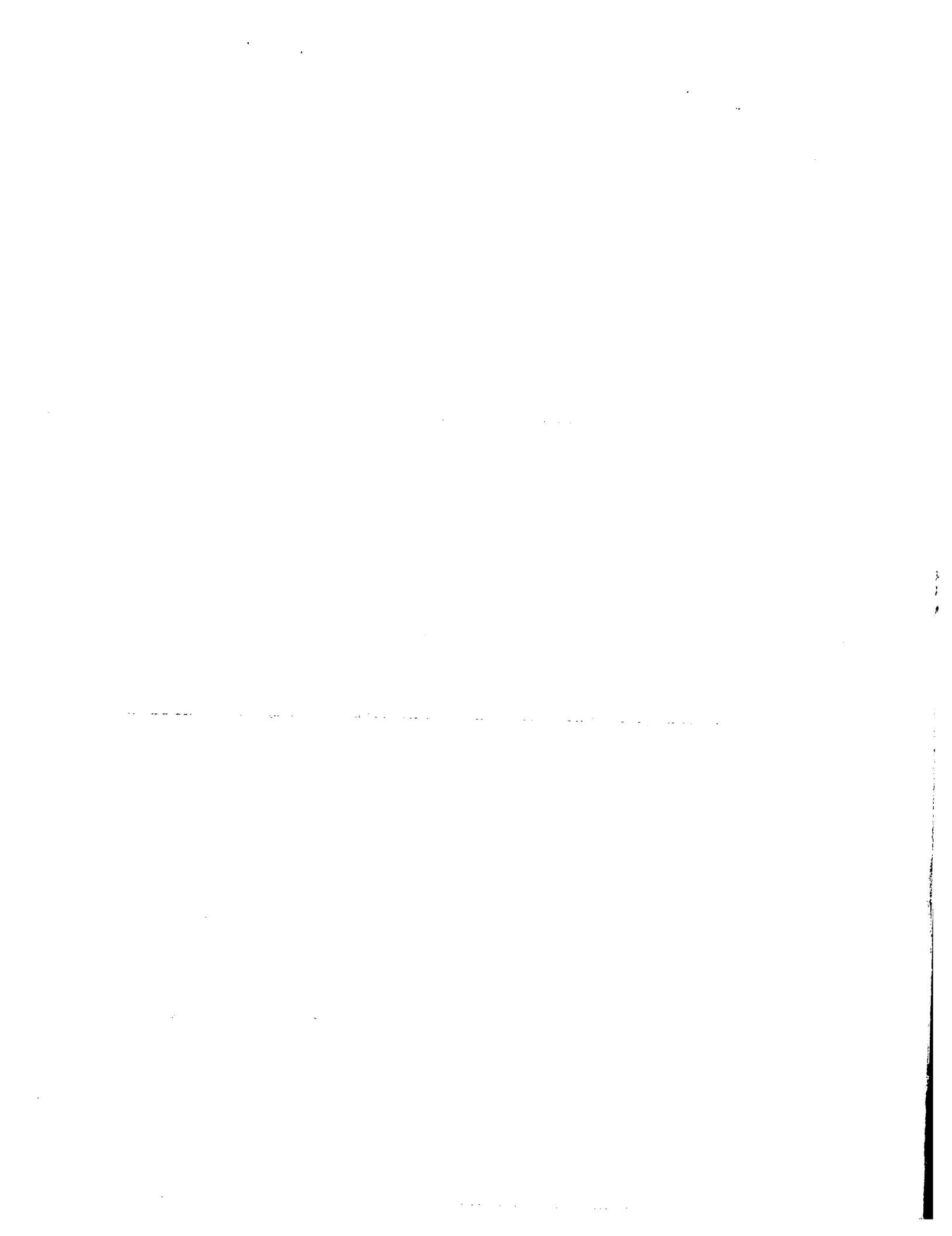
In Anspruch genommene Priorität(en) / Priority(ies) claimed /Priorité(s)
revendiquée(s)
Staat/Tag/Aktenzeichen/State/Date/File no./Pays/Date/Numéro de dépôt:

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PT SE SI SK TR



CATALYST PREPARATION

The present invention relates to the preparation of an epoxidation catalyst and to the process of preparing alkylene oxide with the help of such catalyst.

Background of the Invention

5 An epoxidation catalyst is understood to be a catalyst which catalyses the manufacture of an epoxy group containing compound. A well known process comprises contacting a hydroperoxide and alkene with a heterogeneous epoxidation catalyst and withdrawing a product stream comprising alkylene oxide and an alcohol.

10 Catalysts for the manufacture of an epoxy group containing compound, are well known. EP-A-345856 describes the preparation of such catalyst comprising titanium in chemical combination with a solid silica and/or inorganic silicate. The preparation comprises (a) 15 impregnating a silicium compound with a stream of gaseous titanium tetrachloride preferably comprising an inert gas, (b) calcining the obtained reaction product of step (a), and (c) hydrolysis of the product of step (b). The 20 stream of inert gas also has the function of a carrier for the gaseous titanium tetrachloride. For such use, the gas is to be present in a relatively large amount.

25 There is a continuous interest in improving the selectivity of epoxidation processes in general, and more specifically of processes for the preparation of alkylene oxide. We found a simple and attractive way to achieve this.

Summary of the Invention

The present invention relates to a process for the preparation of an epoxidation catalyst, which process comprises impregnating a silicon containing carrier with a gas stream consisting of titanium halide.

A catalyst of improved selectivity was obtained even though the carrier had been in contact with the same amount of titanium halide.

Detailed description of the invention

10 The catalyst of the present invention is obtained by impregnating a silicon containing carrier. In principle, any silicon containing carrier is suitable for use in the preparation process according to the present invention. Examples of silicon containing carriers comprise
15 zeolites. Preferably, the silicon containing carrier is a silica carrier.

Silica carriers will substantially consist of silicon dioxide. However, limited amounts of further compounds such as contaminants can be present as well.

20 It is known that contaminants can influence the performance of the final catalyst. The silica carrier for use in the present invention preferably contains at most 1200 ppm of sodium, more specifically at most 1000 ppm of sodium. Further, the silica carrier preferably comprises at most 500 ppm of aluminium, at most 500 ppm of calcium, at most 200 ppm of potassium, at most 100 ppm of magnesium and at most 100 ppm of iron. The amounts are based on amount of carrier.

30 The silica carrier preferably is a silica gel. The silica gel carrier for use in the present invention can in principle be any carrier derived from a silicon containing gel. In general, silica gels are a solid, amorphous form of hydrous silicon dioxide distinguished

from other hydrous silicon dioxides by their micro-porosity and hydroxylated surface. Silica gels usually contain three-dimensional networks of aggregated silica particles of colloidal dimensions. They are typically prepared by acidifying an aqueous sodium silicate solution to a pH of less than 11 by combining it with a strong mineral acid. The acidification causes the formation of monosilicic acid (Si(OH)_4), which polymerizes into particles with internal siloxane linkages and external silanol groups. At a certain pH the polymer particles aggregate, thereby forming chains and ultimately gel networks. Silicate concentration, temperature, pH and the addition of coagulants affect gelling time and final gel characteristics such as density, strength, hardness, surface area and pore volume. The resulting hydrogel is typically washed free of electrolytes, dried and activated. A suitable silica gel carrier would be silica support V432 and DAVICAT P-732, which are commercially available from Grace Davison.

Silica gel carriers for use in the present invention preferably have a weight average particle size of at most 2 millimetres. Particle sizes which were found to be especially suitable were weight average particle sizes of from 0.2 to 1.8 mm, more specifically of from 0.4 to 1.6 mm, most specifically of from 0.6 to 1.4 mm.

The silicon containing carrier preferably has a low water content when contacted with the titanium halide. A low water content can be achieved in any way known to someone skilled in the art. A preferred way comprises drying the silicon containing carrier before impregnating the silicon containing carrier with the gas stream consisting of titanium halide. A suitable drying method

comprises subjecting the silicon containing carrier to a temperature of from 200 to 700 °C. Surprisingly it has been found that drying under specific circumstances gives a further improved catalyst. The preferred drying
5 conditions comprise drying the carrier at a temperature of from more than 200 to 300 °C. The drying is preferably carried out during of from 1 to 8 hours, preferably in the presence of an inert gas such as nitrogen. The preferred method has been described in more detail in co-pending patent application claiming priority of European
10 application 02252551.3 (our case TS 1309).

A good performance also was observed if the silicon containing carrier had been subjected to a pretreatment comprising calcining the silicon containing carrier and subsequently hydrolysing the carrier obtained. Hydrolysis comprises treating the carrier with water or steam.
15 Preferably, the hydrolysis is carried out with steam. Alternatively, the hydrolysis treatment may comprise a washing treatment using an aqueous solution of a mineral acid, an aqueous solution of an ammonium salt or a combination thereof. Any water which might still be present after the hydrolysis, is preferably removed before treating the carrier further. Water is preferably removed by drying. Preferably, the calcination is carried
20 out at a relatively high temperature. The preferred high temperature calcination and hydrolysis have been described in more detail in co-pending patent application claiming priority of European application 02252551.3 (our case TS 1062). The treatment comprises (a) calcining a
25 silica gel carrier at a temperature of at least 650 °C,
(b) hydrolysing the calcined silica gel carrier,
(c) impregnating the hydrolysed carrier obtained in
30

step (b) with a titanium-containing impregnating agent, and (d) calcining the impregnated carrier.

The silica gel carrier for use in the present invention preferably has a surface area of at most 5 1000 m²/gram, more preferably at most 800 m²/gram, most preferably at most 500 m²/gram.

Titanium halides which can be used in the process according to the present invention comprise tri- and tetra-substituted titanium complexes which have of from 10 1 to 4 halide substituents with the remainder of the substituents, if any, being alkoxide or amino groups. The titanium halide can be either a single titanium halide compound or can be a mixture of titanium halide compounds. Preferably, the titanium halide comprises at 15 least 50 %wt of titanium tetrachloride, more specifically at least 70 %wt of titanium tetrachloride. Most preferably, the gas stream consists of titanium tetrachloride.

The present invention comprises impregnating the 20 carrier with gas consisting of titanium halide. Surprisingly, it was found that catalyst having higher selectivity for the desired alkylene oxide could be obtained if the silicon containing carrier was 25 impregnated with gas consisting of titanium halide. The preparation according to the present invention is carried out in the absence of a carrier gas. Without wishing to be bound to any theory, it is thought that the carrier gas interferes with the impregnation. However, limited amounts of further gaseous compounds are allowed to be 30 present during the contact between the silicon containing carrier and the gaseous titanium halide. The gas in contact with the carrier during impregnation preferably consists for at least 70 %wt of titanium halide, more

specifically at least 80 %wt, more specifically at least 90 %wt, most specifically at least 95 %wt.

5 Gaseous titanium halide can be prepared in any way known to someone skilled in the art. A simple and easy way comprises heating a vessel containing titanium halide to such temperature that the gaseous titanium halide is obtained.

10 Generally, the impregnated carrier will be calcined and subsequently hydrolysed before being used as a catalyst. It is believed that calcination removes hydrogen halide, more specifically hydrogen chloride which is formed upon reaction of titanium halide and silicon compounds present on the surface of the silicon containing carrier.

15 The optional calcination of the impregnated carrier generally comprises subjecting the impregnated carrier to a temperature of at least 500 °C, more specifically at least 600 °C. Preferably, the calcination is carried out at a temperature of at least 650 °C. From a practical 20 point of view, it is preferred that the calcination temperature applied is at most 1000 °C.

25 Hydrolysis of the impregnated and calcined carrier can remove Ti-halide bonds. The hydrolysis of the impregnated carrier will generally somewhat more severe than the optional hydrolysis of the carrier before impregnation. Accordingly, this hydrolysis of the impregnated carrier is suitably carried out with steam at a temperature in the range of from 150 to 400 °C.

30 Preferably, the hydrolysed impregnated carrier is subsequently silylated for instance by contacting the hydrolysed impregnated carrier with a silylating agent, preferably at a temperature of between 100 and 425 °C. Suitable silylating agents include organosilanes like

tetra-substituted silanes with C₁-C₃ hydrocarbyl substituents. A very suitable silylating agent is hexamethyldisilazane. Examples of specific suitable silylating methods and silylating agents are, for instance, described in US-A-3,829,392 and US-3,923,843 which are referred to in US-A-6,011,162, and in EP-A-734764.

The amount of titanium (as metallic titanium) will normally be in the range of from 0.1 to 10% by weight, suitably of from 1 to 5% by weight, based on total weight of the catalyst. Preferably, titanium or a titanium compound, such as a salt or an oxide, is the only metal and/or metal compound present.

As mentioned above, it is well known in the art to produce alkylene oxides, such as propylene oxide, by epoxidation of the corresponding olefin using a hydroperoxide such as hydrogen peroxide or an organic hydroperoxide as the source of oxygen. The hydroperoxide can be hydrogen peroxide or any organic hydroperoxide such as tert-butyl hydroperoxide, cumene hydroperoxide and ethylbenzene hydroperoxide. The alkene will generally be propene which gives as alkylene oxide, propylene oxide. The catalyst prepared according to the present invention has been found to give especially good results in such process. Therefore, the present invention further relates to a process for the preparation of alkylene oxide which process comprises contacting a hydroperoxide and alkene with a heterogeneous epoxidation catalyst and withdrawing a product stream comprising alkylene oxide and an alcohol and/or water, in which process the catalyst has been prepared according to the present invention.

A specific organic hydroperoxide is ethylbenzene hydroperoxide, in which case the alcohol obtained is 1-phenyl ethanol. The 1-phenylethanol usually is converted further by dehydration to obtain styrene.

5 Another method for producing propylene oxide is the co-production of propylene oxide and methyl tert-butyl ether (MTBE) starting from isobutane and propene. This process is well known in the art and involves similar reaction steps as the styrene/propylene oxide production
10 process described in the previous paragraph. In the epoxidation step tert-butyl hydroperoxide is reacted with propene forming propylene oxide and tert-butanol. Tert-butanol is subsequently etherified into MTBE.

15 A further method comprises the manufacture of propylene oxide with the help of cumene. In this process, cumene is reacted with oxygen or air to form cumene hydroperoxide. Cumene hydroperoxide thus obtained is reacted with propene in the presence of an epoxidation catalyst to yield propylene oxide and 2-phenyl propanol.
20 The latter can be converted into cumene with the help of a heterogeneous catalyst and hydrogen. Suitable processes are described for example in WO 02/48126.

25 The conditions for the epoxidation reaction according to the present invention are those conventionally applied. For propene epoxidation reactions with the help of ethylbenzene hydroperoxide, typical reaction conditions include temperatures of 50 to 140 °C, suitably 75 to 125 °C, and pressures up to 80 bar with the reaction medium being in the liquid phase.

30 The invention is further illustrated by the following Examples.

Examples

The silica gel carrier used in the examples had a surface area of 300 m²/g and a weight average particle size of about 1 mm. Substantially all particles had a
5 particle size between 0.6 and 1.4 mm.

75 grams of silica gel carrier was dried at 250 °C during 2 hours.

The dried silica gel carrier was contacted with a gas stream containing titaniumtetrachloride. The gas stream
10 was obtained by heating titaniumtetrachloride to 200 °C with the help of an electrical heating system. Different gas streams were obtained by adding different amounts of nitrogen. At the end of each experiment, each silica carrier had been in contact with the same amount of
15 titaniumtetrachloride.

The impregnated catalysts thus obtained were calcined at 600 °C during 7 hours. The calcined catalysts were subsequently contacted with steam at 325 °C during
20 6 hours. The steam flow consisted of 3 grams of water per hour and 8 Nl of nitrogen per hour. Finally, the catalysts were silylated at 185 °C during 2 hours by being contacted with 18 grams of hexamethyldisilazane per hour in a nitrogen flow of 1.4 Nl per hour.

The catalysts obtained were analysed for the amount
25 of titanium deposited on the carrier.

The selectivity of the catalysts was tested in a continuous epoxidation bench scale unit containing a number of vessels on automatic weight balances containing respectively the ethylbenzene hydroperoxide and propene
30 feed streams, two high pressure pumps, a fixed bed reactor, a third pump for pumping a recycle stream over the reactor, means to maintain the reactor continuously at temperatures between 60 and 120 °C, a stripper to

remove light boiling components like propene, a cooler and a vessel for receiving the product.

5 The feeds were supplied to the reactor via the two high pressure pumps and mixed together before entering the reactor. The reactor was operated liquid full at 40 bara pressure and 90 °C. A large recycle stream was maintained over the reactor to have isothermal operation of the reactor bed and to ensure that the catalyst to be re-activated is contacted with epoxidation reaction
10 product. The feed was mixed with the recycle stream prior to introduction into the reactor.

The feed consisted of 40 %wt of propene, 20 %wt of ethylbenzene hydroperoxide and 40 %wt of ethylbenzene.

15 The results obtained are given in Table 1. The selectivity is the molar ratio of propylene oxide formed to ethylbenzene hydroperoxide converted.

Example 2

20 Further catalysts were prepared in a way similar to the one described in Example 1. However, the impregnated catalysts were calcined for 6 hours (instead of 7 hours) while the steam flow during the subsequent hydrolysis contained 5 grams of water per hour (instead of 3 grams of water per hour). The results of these experiments are shown in Table 2.

TABLE 1

	Example 1	Comparative Example 1	Comparative Example 2
Impregnating gas %wt titaniumtetrachloride	100	75	60
Catalyst			
titanium (%wt)	4.1	4.0	4.1
sodium (%wt)	0.07	0.07	0.07
Selectivity to propylene oxide			
100-200 hours on stream	90.9	89.7	90.1
200-300 hours on stream	91.3	90.4	91.1

TABLE 2

		Example 2	Comparative Example 3	Comparative Example 4
Impregnating gas	%wt titaniumtetrachloride	100	26	1
Catalyst	titanium (%wt)	3.8	4.1	4.1
	sodium (%wt)	0.07	0.05	0.07
Selectivity to propylene oxide	100-200 hours on stream	90.5	89.6	88.4
	200-270 hours on stream	90.5	89.3	88.3
	270-340 hours on stream	92.5	91.2	90.6
	370-420 hours on stream	92.5	91.4	90.8

C L A I M S

1. Process for the preparation of an epoxidation catalyst, which process comprises impregnating a silicon containing carrier with a gas stream consisting of titanium halide.
- 5 2. Process according to claim 1, in which process the silicon containing carrier is a silica gel.
3. Process according to any one of the preceding claims, in which process the gas stream consists of titanium tetrachloride.
- 10 4. Process according to any one of the preceding claims, in which process the silicon containing carrier comprises at most 1200 ppm of sodium, based on amount of carrier.
5. Process according to any one of the preceding claims, which process comprises drying the silicon containing carrier before impregnation.
- 15 6. Process according to any one of the preceding claims in which process the impregnated carrier is calcined and subsequently hydrolysed.
7. Process according to claim 6, in which process the hydrolysed impregnated carrier is subsequently contacted with a silylating agent.
- 20 8. Process for the preparation of alkylene oxide which process comprises contacting a hydroperoxide and alkene with a heterogeneous epoxidation catalyst and withdrawing a product stream comprising alkylene oxide and an alcohol and/or water, in which process the catalyst has been prepared by a process according to any one of the preceding claims.
- 25

9. Process according to claim 8, in which process the alkene is propene and the alkylene oxide is propylene oxide.
10. Process according to claim 9, in which process the organic hydroperoxide is ethylbenzene hydroperoxide and in which the alcohol is 1-phenyl ethanol.
11. Process according to claim 10, which process further comprises dehydration of 1-phenylethanol to obtain styrene.

A B S T R A C T

CATALYST PREPARATION

Process for the preparation of an epoxidation catalyst, which process comprises impregnating a silicon containing carrier with a gas stream consisting of titanium halide, and process for the preparation of alkylene oxide with the help of such catalyst.

